APR 0 1 2002

OIPE

TECH CENTER 1600/2900

RAW SEQUENCE LISTING

DATE: 12/19/2001

PATENT APPLICATION: US/09/922,067

TIME: 12:34:19

Input Set : N:\Crf3\RULE60\09922067.txt Output Set: N:\CRF3\12192001\I922067.raw

SEQUENCE LISTING

```
3 (1) GENERAL INFORMATION:
            (i) APPLICANT: MacPhee, Colin Houston
     5
                            Tew, David Graham
     6
                            Southan, Christopher Donald
     7
                            Hickey, Dierdre Mary Bernadette
     8
                            Gloger, Israel Simon
     9
                            Lawrence, Geoffrey Mark Prouse
    10
                            Rice, Simon Quentyn John
    11
            (ii) TITLE OF INVENTION: Lipoprotein Associated Phospholipase A2, Inhibitors
    13
                                     Thereof And Use Of The Same In Diagnosis And Therapy
    14
           (iii) NUMBER OF SEQUENCES: 11
    16
            (iv) CORRESPONDENCE ADDRESS:
    18
                  (A) ADDRESSEE: SmithKline Beecham Corporation
    19
                  (B) STREET: 709 Swedeland Road
    20
                  (C) CITY: King of Prussia
    21
                                                              ENTERED
                  (D) STATE: PA
     22
                  (E) COUNTRY: USA
     23
                  (F) ZIP: 19406
     24
             (V) COMPUTER READABLE FORM:
     26
                  (A) MEDIUM TYPE: Diskette
     27
                  (B) COMPUTER: IBM PC compatible
     28
                  (C) OPERATING SYSTEM: PC-DOS/MS-DOS
     29
                  (D) SOFTWARE: PatentIn Release #1.0, Version #1.25 (EPO)
     30
            (vi) CURRENT APPLICATION DATA:
     32
                  (A) APPLICATION NUMBER: US/09/922,067
C--> 33
                  (B) FILING DATE: 03-Aug-2001
C-->34
                  (C) CLASSIFICATION:
     35
           (vii) PRIOR APPLICATION DATA:
     37
                  (A) APPLICATION NUMBER: 09/193,130
     38
                  (B) FILING DATE: 1998-11-17
     39
                  (A) APPLICATION NUMBER: PCT/GB94/01374
     41
                  (B) FILING DATE: 24 June 1994
     42
          (viii) ATTORNEY/AGENT INFORMATION:
     44
                  (A) NAME: Dustman, Wayne J.
     45
                  (B) REGISTRATION NUMBER: 33,870
     46
                  (C) REFERENCE/DOCKET NUMBER: P30693
     47
            (ix) TELECOMMUNICATION INFORMATION:
     49
                   (A) TELEPHONE: 610-270-5023
     50
                  (B) TELEFAX: 610-270-5090
     51
                  (C) TELEX:
     54 (2) INFORMATION FOR SEQ ID NO: 1:
             (i) SEQUENCE CHARACTERISTICS:
     56
                   (A) LENGTH: 37 amino acids
     57
                   (B) TYPE: amino acid
     58
                   (C) STRANDEDNESS:
     59
```

60

(D) TOPOLOGY: linear

1 450 2 01 .

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(ii) MOLECULE TYPE: peptide
     62
           (iii) HYPOTHETICAL: NO
     64
             (v) FRAGMENT TYPE: internal
c--> 68
    70
            (vi) ORIGINAL SOURCE:
c--> 72
            (ix) FEATURE:
            (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:
    74
             Met Leu Lys Leu Lys Gly Asp Ile Asp Ser Asn Ala Ala Ile Asp Leu
     76
                                                  10
     77
             Ser Asn Lys Ala Ser Leu Ala Phe Leu Gln Lys His Leu Gly Leu His
     79
     80
                         20
             Lys Asp Phe Asp Gln
     82
                     35
     83
     85 (2) INFORMATION FOR SEQ ID NO: 2:
             (i) SEQUENCE CHARACTERISTICS:
     87
                  (A) LENGTH: 30 amino acids
     88
     89
                  (B) TYPE: amino acid
                  (D) TOPOLOGY: linear
     90
            (ii) MOLECULE TYPE: peptide
     92
     94
           (iii) HYPOTHETICAL: NO
             (v) FRAGMENT TYPE: internal
     96
             (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:
     100
              Trp Met Phe Pro Leu Gly Asp Glu Val Tyr Ser Arg Ile Pro Gln Pro
     102
                                                    10
     103
              Leu Phe Phe Ile Asn Ser Glu Tyr Phe Gln Tyr Pro Ala Asn
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                           20
     106
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                    (A) LENGTH: 27 amino acids
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                    (B) TYPE: amino acid
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                    (D) TOPOLOGY: linear
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             (ii) MOLECULE TYPE: peptide
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             (iii) HYPOTHETICAL: NO
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              (v) FRAGMENT TYPE: internal
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     123
              Gln Tyr Ile Asn Pro Ala Val Met Ile Thr Ile Arg Gly Ser Val His
     125
     126
              Gln Asn Phe Ala Asp Phe Thr Phe Ala Thr Gly
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                           20
     129
     131 (2) INFORMATION FOR SEQ ID NO: 4:
              (i) SEQUENCE CHARACTERISTICS:
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                    (A) LENGTH: 19 amino acids
     134
                    (B) TYPE: amino acid
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                    (D) TOPOLOGY: linear
     136
             (ii) MOLECULE TYPE: peptide
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             (iii) HYPOTHETICAL: NO
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               (v) FRAGMENT TYPE: internal
     142
              (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:
     146
              Trp Leu Met Gly Asn Ile Leu Arg Leu Leu Phe Gly Ser Met Thr Thr
     148
                                                    10
     149
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1 450 5 01

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	151	Pro Ala Asn												
		(2) INFORMATION FOR SEQ ID NO: 5:												
	156	· · · · · · · · · · · · · · · · · · ·												
	157	57 (A) LENGTH: 420 base pairs 58 (B) TYPE: nucleic acid												
	158													
	159 (C) STRANDEDNESS: double													
	160 (D) TOPOLOGY: linear													
	162													
	164	(iii) HYPOTHETICAL: NO												
c>														
	170													
	172	AAAAAACCTA TTTTAATCCT AATTGTATTT CTCTATTCCT GAAGAGTTCT GTAACATGAT	60											
	174	GTGTTGATTG GTTGTGTTAA TGTTGGTCCC TGGAATAAGA TTCTCATCAT CTCCTTCAAT	120											
*	176	CAAGCAGTCC CACTGATCAA AATCTTTATG AAGTCCTAAA TGCTTTTGTA AGAATGCTAA 180												
	178	TGAAGCTTTG TTGCTAAGAT CAATAGCTGC ATTTGAATCT ATGTCTCCCT TTAATTTGAG	240											
	180	CATGTGTCCA ATTATTTTGC CAGTNGCAAA AGTGAAGTCA GCAAAATTCT GGTGGACTGA	300											
	182	ACCCCTGATT GTAATCATCT TTCTTTCTTT ATCAGGTGAG TAGCATTTTT TCATTTTTAT	360											
	184	GATATTAGCA GGATATTGGA AATATTCAGN GTTGNTAAAA AGNGGNGGCT GAGGGATTCT	420											
187 (2) INFORMATION FOR SEQ ID NO: 6:														
	189 (i) SEQUENCE CHARACTERISTICS:													
	190													
	191													
	192													
	193	(D) TOPOLOGY: linear												
	195													
	197	(iii) HYPOTHETICAL: NO												
C>	199													
	203													
	205	TGCTAATATC ATAAAAATGA AAAAATGCTA CTCACCTGAT AAAGAAAGAA AGATGATTAC	60											
	207	AATCAGGGGT TCAGTCCACC AGANTTTTGC TGACTTCACT TTTGCAACTG GCAAAATAAT	120											
	209	TGGACACATG CTCAAATTAA AGGGAGACAT AGATTCAAAT GTAGCTATTG ATCTTAGCAA	180											
		CAAAGCTTCA TTAGCATTCT TACAAAAGCA TTTAGGACTT CATAAAGATT TTGTTCAGTG	240											
	213	GGACTGCTTG ATTGAAGGAG ATGATGAGAA TCTTATTCCA GGGACCAACA TTAACACAAC	300											
		CAATTCAACA CATCATGTTT ACAGAACTTC TTCCAGGGAA TAGGAGGAAA TACAATTGGG	360											
		GTTTAAAATA GGTTTTTTT	379											
	219	(2) INFORMATION FOR SEQ ID NO: 7:												
	221													
	222													
	223													
	224	• •												
	225													
		227 (ii) MOLECULE TYPE: cDNA												
	229													
C>														
	235 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:													
	237	GAAGAATGCA TTAGATTTAA AGTTTGATAT GGAACAACTG AAGGACTCTA TTGATAGGGA	60 120											
	239	AAAAATAGCA GTAATTGGAC ATTCTTTTGG TGGAGCAACG GTTATTCAGA CTCTTAGTGA AGATCAGAGA TTCAGATGTG GTATTGCCCT GGATGCATGG ATGTTTCCAC TGGGTGATGA	180											
	241	AGATCAGAGA TTCAGATGTG GTATTGCCCT GGATGCATGG ATGTTTCCAC IGGGIGATGA AGTATATTCC AGAATTCCTC AGCCCCTCTT TTTTATCAAC TCTGAATATT TCCAATATCC	240											
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	245	TGCTAATATC ATAAAANTGG AAAAATGCTA CTCACCTGG												
	247	(2) INFORMATION FOR SEQ ID NO: 8:												
	249	(i) SEQUENCE CHARACTERISTICS:												
	250													
	251	· ·												
À	252	(C) STRANDEDNESS: double												
	253													
	255	, ,												
	257													
c>	259	(iv) ANTI-SENSE: NO												
	263	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:												
	265	AAAATAGCAG TAATTGGACA TTCTTTAGGT GGAGCAACGG TTATTCAGAC TCTTAGTGAA												
	267	GATCAGAGAT TCAGATGTGG TATTGCCCTG GATGCATGGA TGTTTCCACT GGGTGATGAA	120											
	269 GTATATTCCA GAATTCCTCA GCCCCTCTTT TTTATCAACT CTGAATATTT CCAATATC 271 GCTAATATCA TAAAAATGAA AAAATGCTAC TCACCTGATA AAGAAAGAAA GATGATTA													
	273 ATCAGGGGTT CAGTCCACCA GAATTTTGCT GACTTCACTT TTGCAACTGG CAAAAT													
	275 GGACACATGC TCAAATTAAA GGGAGACATA GATTCAAATG TAGCTATTGA TCTTAGCAAC													
	277	AAAGCTTCAT CAGCATTCTT ACAAAAGCAT TTAGGACTTC ATAAAGATTT TGATCAGTGG	420											
	279	9 GACTGCTTGA TTGAAGGAGA TGATGAGAAT CTTATTCCAG GGACCAACAT TAACACAACC												
	281 AATCAACACA TCATGTTACA GAACTCTTCA GGAATAGAGA AATACAATTA GGATTAAAAT													
	283 AGGTTTTTA AAAAAAAAA AAAAAAACT CG													
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	287 (i) SEQUENCE CHARACTERISTICS:													
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	289	(B) TYPE: nucleic acid												
	290 (C) STRANDEDNESS: double													
	291	(D) TOPOLOGY: linear												
	293	(ii) MOLECULE TYPE: cDNA												
	295	(iii) HYPOTHETICAL: NO												
C>	297	(iv) ANTI-SENSE: NO												
	300	(ix) FEATURE:												
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	302	, <i>,</i>												
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	308	_ <u>_</u>												
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		CAT GTG CTT TTC TGC CTC TGC GGC TGC CTG GCT GTG GTT TAT CCT TTT	103											
		His Val Leu Phe Cys Leu Cys Gly Cys Leu Ala Val Val Tyr Pro Phe												
	313	10 15 20												
		GAC TGG CAA TAC ATA AAT CCT GTT GCC CAT ATG AAA TCA TCA GCA TGG	151											
		Asp Trp Gln Tyr Ile Asn Pro Val Ala His Met Lys Ser Ser Ala Trp												
	317		100											
		GTC AAC AAA ATA CAA GTA CTG ATG GCT GCT GCA AGC TTT GGC CAA ACT	199											
		Val Asn Lys Ile Gln Val Leu Met Ala Ala Ser Phe Gly Gln Thr												
	321		247											
		AAA ATC CCC CGG GGA AAT GGG CCT TAT TCC GTT GGT TGT ACA GAC TTA	247											
		Lys Ile Pro Arg Gly Asn Gly Pro Tyr Ser Val Gly Cys Thr Asp Leu												
	325	55 60 65 70												

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DATE: 12/19/2001 PATENT APPLICATION: US/09/922,067 TIME: 12:34:19

327	ATG	TTT	GAT	CAC	ACT	AAT	AAG	GGC	ACC	TTC	TTG	CGT	TTA	TAT	TAT	CCA	295
328	Met	Phe	Asp	His	Thr	Asn	Lys	Gly	Thr	Phe	Leu	Arg	Leu	Tyr	Tyr	Pro	
329					75					80					85		
																GAA	343
		Gln	Asp	Asn	Asp	Arg	Leu	Asp	Thr	Leu	Trp	Ile	Pro	Asn	Lys	Glu	
333				90					95					100			
335	TAT	TTT	TGG	GGT	CTT	AGC	AAA	TTT	CTT	GGA	ACA	CAC	TGG	CTT	ATG	GGC	391
		Phe		Gly	Leu	Ser	Lys	Phe	Leu	Gly	Thr	His	Trp	Leu	Met	Gly	
337			105					110					115				
339	AAC	ATT	TTG	AGG	TTA	CTC	TTT	GGT	TCA	ATG	ACA	ACT	CCT	GCA	AAC	TGG	439
			Leu	Arg	Leu	Leu		Gly	Ser	Met	Thr		Pro	Ala	Asn	\mathtt{Trp}	
341		120					125					130					
343	AAT	TCC	CCT	CTG	AGG	CCT	GGT	GAA	AAA	TAT	CCA	CTT	GTT	GTT	TTT	TCT	487
		ser	Pro	Leu	Arg		GLY	Glu	Lys	Tyr		Leu	Val	Val	Phe	Ser	
	135					140					145					150	
34/	CAT	GGT	CTT	GGG	GCA	TTC	AGG	ACA	CTT	TAT	TCT	GCT	ATT	GGC	ATT	GAC	535
	HlS	GIŸ	Leu	GLY		Phe	Arg	Thr	Leu		Ser	Ala	Ile	Gly	Ile	Asp	
349	ama.	001		a	155					160					165		
321	CTG	GCA	TCT	CAT	GGG	TTT	ATA	GTT	GCT	GCT	GTA	GAA	CAC	AGA	GAT	AGA	583
	ьeu	Ата	ser		GLY	Phe	TTe	Va⊥	Ala	Ala	Val	Glu	His	_	Asp	Arg	
353	mam			170					175					180			
355	TCT	GCA	TCT	GCA	ACT	TAC	TAT	TTC	AAG	GAC	CAA	TCT	GCT	GCA	GAA	ATA	631
357	ser	Ald		Ата	Thr	туг	туr		Lys	Asp	GIn	Ser		Ala	Glu	Ile	
	ccc	CAC	185	mom	шаа	ama	m. c	190					195				
320	C111	AGAC	AAG	Com	TGG	CTC	TAC	CTT	AGA	ACC	CTG	AAA	CAA	GAG	GAG	GAG	679
361	GIY	200	пуъ	ser	тъ	теп		ьeu	Arg	Thr	Leu		GIn	GLu	GLu	GIu	
	አሮአ		አጥአ	CCX	74.70	CAC	205	CITIA	000	G33	303	210					
364	Thr	Uic	TIA	Ara	WWI	Clu	CAG	Unl	CGG Arg	CAA	AGA	GCA	AAA	GAA	TGT	TCC	727
365	215	1113	116	ni y	ASII	220	GIII	Val	Arg	GIII		нта	ьys	GIU	Cys		
		GCT	СТС	ልረጥ	СТС		CTT	CAC	ATT	Cam	225	CCA	220	001	ama	230	775
368	Gln	Δla	T.AII	Sor	LAII	Tlo	LAU	Acn	Ile	GWI	Uia	Clrr	AAG	Dma	GTG	AAG	775
369	0.1.1.		Dea	DCI	235	110	neu	дэр	116	240	птэ	GLY	гуѕ	PIO	245	гуѕ	
	ААТ	GCA	מידים	СДТ		ΔAG	ጥጥጥ	САТ	ATG		CAA	CTC	220	CNC		» mm	003
372	Asn	Ala	Leu	Asp	Len	Lvs	Phe	Asn	Met	GIII	Gln	Lau	Lare	Acn	Cor	TIA	823
373				250		110		1155	255	Giu	Q1II	Deu	пуз	260	Set	116	
375	GAT	AGG	GAA		ATA	GCA	GTA	АТТ	GGA	САТ	ጥርጥ	արարա	GGT		CCA	ACC	871
376	Asp	Arq	Glu	Lvs	Ile	Ala	Val	Tle	Gly	His	Ser	Phe	Glv	Glv	Δla	Thr	071
377	-	-	265					270	3 _1		~~	1 110	275	CLY	AIU	1111	
379	GTT	ATT	CAG	ACT	CTT	AGT	GAA		ÇAG	AGA	TTC	AGA	TGT	GGT	ΔͲͲ	GCC	919
380	Val	Ile	Gln	Thr	Leu	Ser	Glu	Asp	Gln	Ara	Phe	Ara	Cvs	Glv	Tle	Ala	212
381		280					285			5		290	010	011		1114	
383	CTG	GAT	GCA	TGG	ATG	TTT	CCA	CTG	GGT	GAT	GAA		тат	TCC	AGA	Δጥጥ	967
384	Leu	Asp	Ala	Trp	Met	Phe	Pro	Leu	Gly	Asp	Glu	Val	Tvr	Ser	Ara	Tle	307
385	295	_		_		300			-	-	305		-1-		5	310	
387	CCT	CAG	CCC	CTC	TTT		ATC	AAC	TCT	GAA		TTC	CAA	TAT	CCT		1015
388	Pro	Gln	Pro	Leu	Phe	Phe	Ile	Asn	Ser	Glu	Tyr	Phe	Gln	Tyr	Pro	Ala	2025
389					315					320	_				325		
391	AAT	ATC	ATA	AAA	ATG	AAA	AAA	TGC	TAC		CCT	GAT	AAA	GAA		AAG	1063

145000

VERIFICATION SUMMARY

PATENT APPLICATION: US/09/922,067

DATE: 12/19/2001 TIME: 12:34:20

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Output Set: N:\CRF3\12192001\1922067.raw

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L:34 M:220 C: Keyword misspelled or invalid format, [(B) FILING DATE:]
L:66 M:220 C: Keyword misspelled or invalid format, [(v) FRAGMENT TYPE:]
L:72 M:220 C: Keyword misspelled or invalid format, [(ix) FEATURE:]
L:74 M:246 W: Invalid value of Alpha Sequence Header Field, [FEATURE:], SeqNo=1
L:166 M:220 C: Keyword misspelled or invalid format, [(iv) ANTI-SENSE:]
L:199 M:220 C: Keyword misspelled or invalid format, [(iv) ANTI-SENSE:]
L:231 M:220 C: Keyword misspelled or invalid format, [(iv) ANTI-SENSE:]
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L:297 M:220 C: Keyword misspelled or invalid format, [(iv) ANTI-SENSE:]